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# **EUROPEAN PATENT SPECIFICATION**

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- (54) Combination therapy comprising ribavirin and interferon alpha in antiviral treatment naive patients having chronic hepatitis c infection

Kombinationstherapie enthaltend Ribavirin und Interferon alpha bei Patienten mit chronischer Hepatitis C Infektion, die nicht antiviral vorbehandelt sind

Théraple combinatoire conténant de la ribavirine et de l'interferon alpha chez les patients ayant une infection chronique d'hépatite C et n'ayant pas été traités par un agent antiviral

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- (56) References cited: EP-A- 0 707 855

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 POYNARD T. ET AL: "A International randomized trial of interferon alfa-2B and ribavirin (intron A/Rebetol) 48 or 24 weeks vs intron A 48 weeks for first line treatment of chronic hepatitis C" HEPATOLOGY, vol. 28, no. 4 Part 2, October 1998 (1998-10), page 387 XP00211025.

# Remarks:

The file contains technical information submitted after the application was filed and not included in this specification

Description

#### BACKGROUND OF THE INVENTION

Dot11 The present invention relates to uses of ribavinin and interferon alpha to prepare pharmaceutical compositions for treating antiviral treatment naive patient having chronic hepatitis C infection to eradicate detectable HCVANA involving a combination therapy using a therapeutically effective amount of ribavinin and a therapeutically effective amount of interferon-alpha for a time period of from 40 up to 50 weeks. According to the invention the patient is one having a HCV genotype type 1 infection and a vital lead of greater than 2 million copies per ml as measured by HCV-RNA quantitative PCR.

[0002] Chronic infection with hepatitis C virus is an insidious and slow-progressing disease having a significant impact on the quality of life. It can eventually result in cirrhosis of the liver, decompensated liver disease and/or hepatocelluar carchoma.

[0003] Alpha interferon monotherapy is commonly used to treat chronic hepatitis C infection. However, this treatment is not always effective and sometimes results in intolerable side effects related to the dosage and duration of therapy. Ribavirin has been proposed as a monotherapy treatment for chronic hepatitis C infection (Thomas et al. AASLD Abstracts, Hepatology Vol. 20, NO. 4, Pt 2, Number 440, 1994), However, this monotherapy treatment has usually been found ineffective. Combination therapy of alpha interferon and ribavirin has been proposed: (Lai, et al. Symposium to the 9th Biennial Scientific Meeting Asian Pacific Association for the Study of the Liver, 1994); Combination treatment with interferon alpha-2b and ribavirin for chronic hepatitis C in patients who have failed to achieve sustained response to interferon alone: Swedish experience. J Hepatology, 1995;232 (Suppl 2):17-21. Brouwer JT, Nevens F, Michielsen P, et al.; What options are left when hepatitis C does not respond to interferon? Placebo-controlled Benelux multicentre retreatment trial on ribavirin monotherapy versus combination with interferon. J Hepatol. 1994;212 (Suppl 1):S17. Abstract WP2/08. Chemello L, Cavalletto L, Bernardinello E, et al. Response to ribavirin, to interferon and to a combination of both in patients with chronic hepatitis C and its relation to HCV genotypes. J Hepatol. 1994;212 (Suppl 1):S12. Abstract GS5/29; and The effect of interferon alpha and ribavirin combination therapy in naive patients with chronic hepatitis C, J. Hepatol. 1995;23(Suppl.2):8-12. Reichard et al. LANCET 1998; 351;83-87 disclosed that more chronic hepatitis C patients have a sustained virologic response with a combination of interferon alpha-2b and ribavirin for 24 weeks than with only interferon alpha-2b. Reichard et al. also disclosed that interferon-alpha-2b alone is sufficient to achieve a sustained response in such patients with HCV-RNA serum values above 3 million copies/mL. However, no one has described methods using alpha interferon and ribavirin which eradicate HCV-RNA in any long-term, effective manner for antivirally naive patients having a specific HCV genotype infection.

[0004] There is a definite need for a treatment of antiviral treatment naive patients having chronic hepatitis C infection with a combination of alpha interferon and ribavirin which eradicates HCV-RNA in any long-term, effective manner.

# SUMMARY OF THE INVENTION

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[0005] The invention involves improved uses of ribavirin and interferon alpha to prepare pharmaceutical compositions for treating antiviral treatment naive patient having chronic hepatitis C infection to eradicate detectable HCV-RNA involving a combination therapy using a therapeutically effective amount of ribavirin and a therapeutically effective amount of interferon-alpha for a time period of from 40 up to 50 weeks as defined in the claims.

[0006] We have discovered that if the antiviral treatment naive patient has HCV genotype 1 infection, or if the antiviral treatment naive patient has an HCV genotype 1 infection, and a viral load of greater than 2 million of HCV-RNA by quantitative PCR, then the administration of the combination therapy is effected for a time period of 40-50 weeks, preferably 48 weeks.

[0007] According to the present invention, there is provided

- the use of ribavirin for the manufacture of a pharmaceutical composition for treating a patient having chronic hepatitis C infection to eradicate detectable HCV-RIAN wherein the pharmaceutical composition is for administering an effective amount of ribavirin in association with an effective amount of interferon alpha, characterised in that the ribavirin in association with the interferon alpha is for administration for a time period of 40-50 weeks, the patient is an antiviral treatment naive patient, and the patient is one having a HCV genotype type 1 infection and a viral load of greater than 2 million copies per mI of serum as measured by HCV-RNA quantitative PCR.
  - 5 the use of interferon alpha for the manufacture of a pharmaceutical composition for treating a patient having chronic hepatitis Clinfection to eradicate detectable HCV-RNA wherein the pharmaceutical composition is for administering an effective amount of interferon alpha in association with an effective amount of ribavirin, characterised in that the ribavirin in association with the interferon alpha is for administration for a time period of 40-50 weeks. the

patient is an antiviral treatment naive patient, and the patient is one having a HCV genotype type 1 infection and a viral load of greater than 2 million copies per ml of serum as measured by HCV-RNA quantitative PCR. the use of both ribavinin and interferon alpha for the manufacture of pharmaceutical composition for treating a patient having chronic hepatitis C infection to eracleate detectable HCV-RNA wherein the pharmaceutical composition is or administering an effective amount of ribavirin in association with an effective amount of interferon alpha characterised in that the ribavirin in association with the interferon alpha is for administration for a time period of 40-50 weeks, the patient is an antiviral treatment naive pleatint, and the patient is one having a HCV genotype type 1 infection and a viral load of greater than 2 million copies per ml of serum as measured by HCV-RNA quantitative PCR.

[0008] The interferon-alpha administered is preferably selected from interferon alpha-2a, interferon alpha-2b, a consensus interferon, a purified interferon alpha product or a pegylated interferon alpha-2a or pegylated interferon alpha-2b. More preferably, the interferon-alpha is selected from interferon alpha-2b, or a purified interferon alpha-2b, or a purified interferon alpha-2 product and the amount of interferon-alpha administered is from 2 to 10 million IU per week on a weekly, TIM, QOD or daily basis. In a preferred embodiment, the interferon-alpha administered is interferon-alpha-2b and the amount of interferon-alpha administered is million IU TIM.

[0009] Alternatively, the interferon-alpha administered is consensus interferon and the amount of interferon-alpha administered is from 1 to 20 micrograms per week on a weekly, TIW, OOD or daily basis. In another embodiment, the interferon-alpha administered is a pegylated interferon alpha 2b and the amount of pegylated interferon-alpha 2b administered is from 0.5 to 2.0 micrograms/filogram per week on a weekly, TIW, OOD or daily basis. Alternatively, the interferon-alpha administered is a pegylated interferon alpha-2a and the amount of pegylated interferon-alpha-2b and the amount of pegylated interferon-alpha-2b and the amount of pegylated interferon-alpha-2b and the amount of pegylated interferon-alpha administered is from 20 to 250 micrograms/filogram per week on a weekly, TIW, OOD or daily basis.

[0010] The use of interferon alpha-2a or pegylated interferon alpha-2a or interferon alpha-2b or pegylated interferon alpha-2b is preferred.

[0011] During the 40-50 week time periods, the amount of ribavirin administered is 800 to 1200 mg per day, preferably 800, 1000 or 1200 mg per day, and the amount of interferon alpha-2a or interferon alpha-2b administered is from 2 to 10 million IU TW.

#### DETAILED DESCRIPTION

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[0012] Surprisingly, it has been found that, in the case of antiviral treatment naive patients having chronic hepatitis C infection and having HCV genotype 1, or such naive patients having HCV genotype 1 and a viral load of greater than 2 million copies per ml of HCV-RNA by quantitative PCR ("PCR"), combination therapy with a therapeutically effective amount of ribavirin and a therapeutically effective amount of interferon alpha for a time period of at least 20 to 30 weeks results in ten times more patients having no detectable HCV-RNA in their serum at least 24 weeks after termination of therapy compared to by interferon-alpha monotherapy. When the combination therapy is extended to a time period of 40 to 50 weeks, two to three times more patients have no detectable HCV-RNA in their serum at least 24 weeks after termination of combination therapy compared to those treated with the combination therapy for 24 weeks and eight to nine times more patients have no detectable HCV-RNA in their serum at least 24 weeks after termination of combination therapy compared to those treated with interferon-alpha monotherapy for 48 weeks. See Tables 6, 14, 16 &17, the rate of sustained virologic response found after using the combination therapy depends upon the HCV genotype and the base line viral load as measured by HCV-RNA/qPCR as well as the treatment period of the combination therapy for HCV genotype 1. See Tables 13 & 15. The treatment period of the combination therapy for antiviral treatment naive patients having chronic HCV genotypes 4, 5 and 6 infections is the same as antiviral treatment naive patients having chronic naive patients having chronic HCV genotype 1. The treatment period of the combination therapy for antiviral treatment naive patients having HCV genotypes 2 and/or 3 is shorter, namely 20 to 30 weeks, preferably 24 weeks. See Tables 7, 13 & 15.

19013] The term "Interferon alpha" as used herein means the family of highly homologous species-specific proteins that inhibit vial replication and cellular proliferation and modulate immune response. Typical suitable interferon-alphas include, but are not limited to, recombinant interferon alpha-2s such as Intron-A interferon available from Schering Corporation, Kenilworth, N.J., recombinant interferon alpha-2s such as Roferon interferon available from Hoffmann-La Roche, Nutley, N.J., recombinant interferon alpha-1a, a purified blend of natural alpha interferons such as Sumiferon available from Boeringer Ingelheim Pharmaceutical, Inc., Ridgefield, CT., interferon alpha-11, a purified blend of natural alpha interferons such as Sumiferon available from Sumitonon, Japan or a Welfleron interferon alpha-11 (INS) available from the Glasso-Wellcome Ltd., London, Great Britain, or a consensus alpha interferon such as those described in U.S. Patent Nos. 4,897.471 and 4,995.623 (sepocially Examples 7, 8 or 9 thereof) and the specific product available from Amgen, Inc., Newbury Park, CA, or interferon alpha-3 a mixture of natural alpha interferons made by Interferon Sciences and available from the Purdue Frederick Co., Norwalk, CT., under the Afferon Tadenamen. The use of Interferon alpha-16.

or alpha 2b is preferred. Since interferon alpha 2b, among all interferons, has the broadest approval throughout the world for treating chronic hepatitis C infection, it is most preferred. The manufacture of interferon alpha 2b is described in U.S. Patent No. 4,530,901.

[0014] The interferon alpha administered is selected from interferon alpha-2a, interferon alpha-2b, a consensus interferon, a purified interferon alpha product or a pegylated interferon-alpha-2a or pegylated interferon alpha-2b.

[0015] The therapeutically effective amount of interferon alpha-2a, interferon alpha-2b, or a purified interferon alpha administered in association with ribavirin is from 2 to 10 million IU per week on a weekly, TIW, QOD or daily basis.

[0016] The therapeutically effective amount of interferon-alpha-2b administered is 3 million IU TIW.

[0017] When the interferon alpha administered in association with ribavirin is consensus interferon, the therapeutically effective amount of interferon-alpha administered is from 1 to 20 micrograms per week on a weekly, TIW, QOD or daily basis.

[0018] The term "pegylated interferon alpha" as used herein means polyethylene glycol modified conjugates of interferon alpha, preferably interferon alpha, 2a not alpha. 2b. The preferad polyethylene glycol-interferon alpha 2b not alpha. 2b. The phrases "12,000 molecular weight polyethylene glycol conjugated interferon alpha" and "PEG<sub>12000</sub>; IPN alpha" as used herein mean conjugates such as are prepared according to the methods of international Application No. WO 95/13090 and containing urethane linkages between the interferon alpha-2 or -2b amino groups and polyethylene glycol having an average molecular weight of 12000. The pegylated interferon alpha, PEG<sub>1200</sub>; IFN-alpha-2 is a valiable from Schering-Plough Research institute, Kenliworth, N.

[019] The preferred PEG<sub>1900</sub>-interferon alpha-2b is prepared by attaching a PEG polymer to the epsilon amino group or a lysine residue in the interferon alpha-2b molecule. A single PEG<sub>1900</sub> molecule is conjugated to free amino groups on an IFN alpha-2b molecule via a urethane linkage. This conjugate is characterized by the molecular weight of PEG<sub>1900</sub> and stached. The PEG<sub>1900</sub> PIN alpha-2b conjugate is formulated as a lyophilized powder for injection. The objective of conjugation of interferon alpha with PEG is to improve the delivery of the protein by significantly prolonging its plasma half-life, and thereby provide protracted activity of interferon alpha.

[0020] Other Interferon alpha conjugates can be prepared by coupling an interferon alpha to a water-soluble polymer. A non-limiting list of such polymers include often polyalkylene outdo homopolymers such as polypropylene glycost, polyoxyethylenated polyols, copolymers thereof and block copolymers thereof. As an alternative to polyalkylene outdo-based polymers, effectively non-antiligenic materials such as dextra, polyvinlypyrolidones, polyacryfamides, polyyingl alcohols, carbohydrate-based polymers and the like can be used. Such interferon alpha-polymer conjugates are described in U.S. Patent No. 4,767,68, U.S. Patent No. 4,776,68, U.S. Patent No. 4,776,68, U.S. Patent No. 4,776,88, European Patent Application No. 205 897, European Patent Application No. 205 897, European Patent Application No. 205 897, European Patent No. 517,098, 205 897, European Patent No. 4,776, 205 897, European Patent No. 517,098, 205 897, 205 89

[0021] Pharmaceutical compositions of pegylated interferon alphasuitable for parenteral administration may be formulated with a suitable buffer, e.g., Tris-HOL, cotate or phosphate such as dibacic sodium phosphate/monobasic sodium phosphate buffer, and pharmaceutically acceptable excipients (e.g., sucrose), cerriers (e.g., human plasma albumin), toxicity agents (e.g., Naci), preservatives (e.g., thimenosol, cresol or beny) alcohol), and surfactants (e.g., tween or polysorbates) in sterile water for injection. The pegylated interferon alpha-may be stored as typibilized powders under a refrigeration at 2°-8°C. The reconstituted aqueuous solutions are stable when stored between 2° and 8°C and used within 24 hours of reconstitution. See for example U.S. Patent Nos., 4492,537,5,762,923 and 5,766,582. The reconstituted aqueuous solutions may also be stored in prefilled, multibose syringes such as those useful for delivery of drugs such as insulin. Typical suitable syringes include systems comprising a prefilled year level psyringes with a strength of the psyringe syringes with a strength of the psyringe syringe syringe syringe such as the NOVOLET Novo Pen available from Novo Novilak, as well as prefilled, pen-type syringes withdiallow seasy self-injection by the user. Other syringe systems include a pen-type syringe compartment.

[0022] When the interferon-alpha administered in association with ribavirin is a pegylated interferon alpha-2b and the amount of interferon-alpha administered is from 0.5 to 2.0 micrograms/fillogram per week on a weekly, TTW, QOD or daily basis.

[0023] When the interferon-alpha administered in association with ribavirin is a pegylated interferon alpha-2a and the amount of interferon-alpha administered is from 20 to 250 micrograms/kilogram per week on a weekly, TIW, QOD or daily basis.

[0024] Ribavirin, 1-β-Orthofuranosyi-1H-1,2.4-triazole-3-carboxamide, available from ICN Pharmaceuticals, Inc., Costa Mesa, California, is described in the Merck Index, compound No. 8199, Eleventh Edition. Its manufacture and formulation is described in U.S. Patent No. 4,211,771.

[0025] A person suffering from chronic hepatitis C infection may exhibit one or more of the following signs or symptoms:

(a) elevated ALT.

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- (b) positive test for anti-HCV antibodies.
- (c) presence of HCV as demonstrated by a positive test for HCV-RNA.
- (d) clinical stigmata of chronic liver disease.
  - (e) hepatocelluar damage.

[9026] To practice the invention, the combination therapy of interferon alpha and ribavirin are administered to the patient exhibiting one of more of the above signs or symptoms in amounts sufficient to eliminate or at least alleviate one or more of the signs or symptoms. Interferon alpha formulations, are not effective when administered orally, so the preferred method of administering the interferon alpha formulations, are not effective when administered orally, so the preferably by subcutaneous, IV, or IM, injection. The ribavirin is administered to the patient in association with the interferon alpha, that is, the Interferon alpha dose is administered during the same period of time that the patient receives doses of habvirin, Ribavirin may be administered orally in capsule, tablet or liquid form in association with the parenteral administration of pegylated interferon-alpha. Of course, other types of administration of some medicaments, as they become available are contemplated, such as by neasal spray, transdermally, by suppository, by sustained release dosage form, and by pulmonary inhalation. Any form of administration of loth ropper dosages are delivered without destroying the active ingredient active ingredient and earlier interferon.

0 [0027] The term "antivral treatment naive patients" in the context of the present invention means that the patients have never been treated with ribavirin or any interferon including, but not limited to an interferon-alpha.

[0028] The term "not perfectly of detectable HCV-RNIA" in the context of the present invention means that there is less than 100 copies of HCV-RNIA per mil of serum of the patient as measured by quantitative, multi-cycle reverse transcriptase PCR methodology, HCV-RNIA is preferably a measured by expension by the methodology described below. This methodology services are the present invention by the methodology described below. This methodology is a CV-RNIA by the present invention by the methodology described below.

[0029] RNA is extracted from patient serum using a guaninditum thicocyanate- phenot-chloroform mister followed by ethanol-ammonium acetate precipitation. The precipitated RNA is centrifuged and the resulting pellet is dried in a Centrivap console (Labconco, Karasas City, Mo.). The dry pellet is then resuspended in 30 milrorillers of an Rnasin (Promega Corp., Madison, WI), dithiothritol, and diethlyprocarbonate-treated water mixture. Samples are kept at or below -20°C (preferably below "-0°C (putil RNA everse transcription (RT) and PCR).

[0030] In order to convert the entire RNA sequence into cDNA in the RT reaction, random hexadeoxyribonucleoides (Pharmacia Biotech, Piscatavays, U)) are used as primers for the first stand cDNA synthesis. Two aliquots of 3 microliters of resuspended sample is added to 3 microliters of 100ng/ul random primers and denaturated at 7°CC, then verses transcribed at 40°C for one hour using M-MLY reverse transcriptace (USB, Clevvielan, CH) in standard tuffer containing 5 mM MgCl<sub>2</sub>. The final RT reaction volume is 26 µL The PCR is started immediately following the reverse transcriptace.

[0031] A modified version of the PCR method is performed using heat-stable Taq polymerase to amplify the cDNA. Seventy-five microliters of PCR mix is added to the entire RT reaction volume (26 μl) to a final MgCl<sub>2</sub> concentration of 1.5 mM in a total volume of 101 μl. Each 101 μl sample is then split into 50.5 μl, and a layer of mineral oil is placed on too to prevent evaporation.

[00.32] The PCR cycle consists of annealing for 90 sec, extension for 90 sec, and denaturation for 90 sec, at 55°X, 74°C and 94°C, respectively. Thermocycling samples is submitted to a final 74°C extension for 10 minutes. Four different cycle sets are used. By loading the sample in duplicate, and splitting these samples evently after RT, there are four tubes from one sample. Each of the four tubes is given a different cycle number, enhancing sensitivity and accuracy in the quantitation process. The thermocycling elificancy will be assessed by satisfactory amplification of known copy number RNA standards included in each set of 60 tubes. Two primer sets are used for the amplification, both from the 5° untranslated region of the HCV genome. Both of these primer sets are highly conserved and detect all known subtypes of HCV. Primer set 1: upstream 5°-GTG GTC TGC GGA ACC GGT GAA CT ACT GTC TTC-3′ which produced a 190 bp product. Primer set 2: upstream 5°-GTG TGA GGA ACT ACT GTC TTC-3′ dworstream 5°-CTG TGA GGA ACT ACT GTC TTC-3′ ACM STEAR CAAS within produced a 256 bp product.

[0033] The amplified cDNA is then electrophorised in 3% agarose gel and transferred to nylon membrane. The target DNA is detected by Southern blotting and immunostaining using a nonradioactive dipoxigenin-labeled DNA probe. These procedures are performed using automated instruments for PCR thermocycling, agarose get electrophoresis, vacuum-transfer Southern blot, hyndrization, and immunostaining. Each membrane contains known copy number serially diluted standards which are used to construct standard curves for quantitative measurement of the specimen bands. Originally standard curves are made from carefully diluted HCV-RNA from transcribed clones. Radioactive incorporation studies, get electrophoresis, and OD 260 are performed on the transcripts to determine that they are of the expected length. After the production of the RNA transcribts quantitated clone standards "posiced" standards as

generated which better represent the heterogeneous nature of HCV, one would encounter in natural infection. These pools are made by combining large amounts of serum or plasma from known infected individuals. The serum/plasma pools are calibrated with PCR, against the clone transcripts and then diluted in the known PCR-negative fluids. Finally, the higher copy number samples of the pools are checked against the cDNA Quantiplex nucleic acid detection system from Chiron Inc. (Emeryville, CA). These "double quantitated" pools are aliquoted and saved at -70°C. Dilutions of 5,000,000, 1,000,000, 500,000, 100,000, 10,000, and 1000 copies/ml are used in each experiment.

[0034] Each Southern blot membrane is scanned into a computer using an automated scanner/densitometer, at intervals during development to determine when the standard curve is most linear. The resultant electronic images are then measured for band area and mean band density. All of the reading are standardized to integrated band density and compared to the standard curve to obtain a numerical value of viral copy number for each band.

[0035] The term "sustained virologic response" as used in the context of the present invention means that there is no detectable HCV-RNA in the serum of patients treated in accordance with the present invention for at least 24 weeks after the end of the combined therapy treatment, Preferably, the period of sustained virologic response is at least one year - or longer - after the end of treatment.

[0036] The following clinical protocols were performed:

Study 1:

#### Overall Design and Plan of the Study 20

[0037] This was a prospective, multicenter, randomized, double-blind, parallel-group. The study compared treatment with INTRON® A plus ribayirin to treatment with INTRON® A plus placebo for 24 or 48 weeks in antiviral treatment naive patients with compensated chronic hepatitis C who had no prior treatment with any interferon including but not limited to alpha interferon (INTRON® A. Roferon®-A. consensus interferon, or Wellferon®) therapy and who also had no prior treatment with ribavirin. Patients who had prior treatment for hepatitis with any other antiviral or immunomodulatory drug within the previous 2 years were also excluded from this study. Eligible patients had chronic hepatitis C confirmed by positive serum HCV-RNA, liver biopsy, and laboratory tests,

[0038] Patients were randomized to treatment with either INTRON® A plus ribavirin or INTRON® A plus placebo. The dose of INTRON® A was 3 million IU SC TIW; the dose of ribavirin was 1000 or 1200 mg PO daily (based on weight) in two divided doses. Treatment group assignments were made in equal ratios by a Central Randomization Center. The randomization procedure was designed to attempt to balance the treatment groups, within and across sites, with respect to presence or absence of cirrhosis in the pretreatment liver biopsy, serum HCV-RNA/qPCR level, and HCV genotype.

[0039] Study treatment was administered for 24 or 48 weeks. The total course of the study was 48 or 72 weeks to determine long-term effect of treatment. Duration of treatment was assigned at the time of randomization.

[0040] During treatment and posttreatment follow-up, biochemical (ALT), virological (HCV-RNA), and histological (liver biopsy) examinations were used to assess the nature and duration of response to study treatment. The primary efficacy variable was the overall response defined as loss of serum HCV-RNA/qPCR (<100 copies/mL) as measured at24 weeks following the end of therapy. In addition, a decrease in hepatic inflammation, an I mprovement in posttreatment liver biopsy as measured by the Knodell Histology Activity index (HAI) and normalization of ALT were also examined as a secondary efficacy endpoints. The safety of the study treatments was assessed by monitoring selected laboratory parameters and by also recording and evaluating the occurrence of any adverse events.

# Treatment Regimens

[0041] There were four study treatment regimens:

- 1. INTRON® A 3 million IU SC TIW plus ribayirin 1000 or 1200 mg/day PO in two divided doses for 24 weeks; or
- 2. INTRON® A 3 million IU SC TIW plus placeho matching ribayirin PO in two divided doses for 24 weeks; or 3. INTRON® A 3 million IU SC TIW plus ribayirin 1000 or 1200 mg/day PO in two divided doses for 48 weeks; or
- - 4. INTRON® A 3 million IU SC TIW plus placebo matching ribavinn PO in two divided doses for 48 weeks.

[0042] Study treatments 1 and 2 were administered for 24 weeks; study treatments 3 and 4 were administered for 48 weeks. The standard INTRON® A (interferon alpha-2b, recombinant) regimen for hepatitis C was administered as a fixed dose of 3 million IU TIW. Each patient received instructions regarding the preparation and subcutaneous administration of INTRON® A. Ribavirin was administered twice daily, morning and evening. The dose was determined by the patient's body weight at the Entry visit. Patients weighing ≤75 kg received 1000 mg daily as two 200 mg capsules in the morning and three 200 mg capsules in the evening. Patients weighing >75 kg received 1200 mg daily as three

200 mg capsules morning and evening.

[0043] The randomization procedure was designed to balance the groups with respect to the following Baseline characteristics:

- pretreatment liver histology (cirrhosis or no cirrhosis);
  - serum HCV-RNA/aPCR status (HCV-RNA/aPCR ≤2.000,000 or HCV-RNA/aPCR >2.000,000 copies/mL); and
  - HCV Genotype (1 or other). Patients with mixed genotypes (which include Type 1) will be classified as Type 1 for purposes of balancing.

# 10 Efficacy

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[0044] The primary efficacy objective was comparison of the freatment groups 1 and 2 and 3 and 4 with respect to the sustained virologic response rate defined as loss of (detectable) serum HCV-RNA/pCRC measured at 24 weeks following the end of therapy to an undetectable level or to a level <100 copies/mL. The following secondary efficacy Endotistive very also examined:

[0045] The secondary efficacy Endpoints:

- · proportions of patients with normalization of ALT at 24 weeks of follow-up;
- proportions of patients with improvement in biopsy (Categories I + II + III combined scores);
  - changes from Baseline in the biopsy scores (Categories I + II + III combined scores);
  - response rates at Endpoint of treatment based on HCV-RNA/aPCR;
  - · proportion of patients with normalization of ALT at Endpoint of treatment.
  - response rates at 24 weeks of follow-up based on HCV-RNA/aPCR.

# 30 Virology: Entry Status and Change from Entry

[0046] Serum HCV-RNA/qPCR testing was performed by a central laboratory. A positive HCV-RNA assay result was required at Baseline; only patients positive for HCV-RNA were eligible to participate. Repeat assays were scheduled at Weeks 4, 12, 24, and if the patient was in the 48 week treatment groups at weeks 38 and 48. All patients had repeat assays scheduled for Follow-up Weeks 12 and 24.

[0047] Response was assessed as defined below:

[0048] Responder: A patient was classified as a responder at a given time point if HCV-RNA/qPCR was negative (<100 copies per mL) at that time point.

[0049] Sustained Responder: A patient was classified as a sustained responder if the patient was a responder at 24 weeks of follow-up.

[0050] Note that patients who do not meet these criteria, including patients who discontinued before the required HCV-RNA/qPCR evaluations were obtained, were classified as non-responders.

[0051] Overall Responder: Based on both serum HCV-RNM<sub>2</sub>PCR and change in liver histology as evaluated by the Knodel HAI Inflammation Score. A patient was classified as an overall responder to treatment if he/she was a sustained responder and his/her Post treatment Knodel HAI inflammation score (sum of categories i+ii+iii) improved by 2 or more units relative to the Pretreatment score.

### Liver Histology

Viol [0052] Liver biopsy was required within the six months preceding patient enrollment and at Follow-up Week 24 for all patients. Evaluation of the biopsies was performed by a single pathologist using the Knodell Histology Activity Score. The central pathologist was blinded with respect to patient identification, renamengroup, and the time the biopsy obtained relative to treatment/Pre-or Posttreatment). Efficacy of study treatments was assessed by comparing the degree of inflammatory activity observed at Baseline with that present at Follow-up Week 24.

#### RESULTS

[0053] Nine hundred-twelve patients were enrolled at 42 US centers and randomized to treatment with either IN-

TRON® A plus ribavirin (N=228) or INTRON® A plus placebo (N=23) for 24 weeks or treatment with either INTRON A plus ribavirin ("I+R") (N=228) or INTRON A plus placebo ("I+P") (N=225) for 48 weeks.

[0054] Overall, 81% (734/912) of patients completed treatment and 24 weeks of follow-up. Eighty-nine percent (2032/28) of patients in the 24 week I+P group, 95% (207/231) of patients in the 24 week I+P group, 75% (159/232) of patients in the 48 week I+P group completed the study. [0055] Twenty percent (178/912) of patients isocontinued during treatment 11% (25/228) in the 24 week I+P group, 10% (24/231) in the 24 week I+P group, 30% (69/228) in the 48 week I+P group, and 27% (60/228) in the 48

[0056] At least 96% of patients who completed treatment and entered follow-up completed the study. Only 2 patients in the 14\* 24 week group, 8 patients in the 24 week 14\* 24 week group, 8 patients in the 24 week 14\* 24 week group, and 4 patients in the 48 week 14\* prorup discontinued durino flow-up.

[0057] The patient's weight and their baseline disease characteristics (HCV genotype and initial viral load) for all patients in Study 1 is given in Table 1 below. HCV genotypes were done on the patient serum samples subjected to HCV-PNAVQPCR testing.

Table 1

Table 1.					
Body Weight and Bas	seline Diseas	e Characteri	stics for Stud	ly 1 Patients	
Weight	I+R <sup>1</sup> 24 Wks (N=228)	I+P <sup>2</sup> 24 Wks (N=231)	I+R1 48 Wks (N=228)	I+P <sup>2</sup> 48 wks (N=225)	
>75 kg <75 kg	148 (65%) 80 (35%)	157 (68%) 74 (32%)	133 (58%) 95 (42%)	153 (685) 72 (32%)	
HCV Genotype <sup>3</sup>					
1 2 3 4 5 6 HCV-RNA/gPCR	164 (72%) 29 (13%) 28 (12%) 6 (3%) 0 1 (0.4%)	167 (72%) 38 (17%)_ 24 (10%) 2 (0.9%) 0	166 (73%) 37 (16%) 23 (10%) 1 (0.4%) 1 (0.4%) 0	162 (72%) 43 (19%) 19 (8%) 1 (0.4%) 0	
(copies/ <sub>mL)</sub> ,					
Geometric Mean ≤ 2 million copies/mL > 2 million copies/mL	3,070,019 62 (27%) 166 (73%)	2,767,469 74 (32%) 157 (68%)	2,922,925 76 (33%) 152 (67%)	2,819,324 63. (28%) 162 (72%)	

# FOOTNOTES TO TABLE 1

1. I + R is Intron A + Ribavirin

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- 2. I + P is Intron A + Placebo
  - Sub-genotypes are classified under their respective genotype.

[0058] All discussions of efficacy and safety in this report are based on data for the all-treated groups.

# Efficacy

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<sup>10</sup> [0059] The objectives of this study were to compare INTRON® A plus rhawrin with INTRON® A plus placebo with respect to the overall response rate and the viologic response rate (based on HCV-RNA (qPCR) for 24 and 48 weeks. The primary efficacy variable for the study is the overall response rate.

[0061] The combination therapy of INTRON A plus ribavirin administered for 48 weeks increased by 2 to 3-fold the efficacy of INTRON A monotherapy for the treatment of chronic hepsilis C in antivirul treatment naive patients. Forty-eight weeks of the combination therapy of INTRON A plus ribavirin increased the response rate at the End of Treatment and decreased the rate of relapse, which resulted in a better sustained viorolgic response rate than for 44 weeks of

INTRON A +Placebo. This enhancement of efficacy included all aspects of the disease and resulted in:

- Sustained eradication of detectable HCV-RNA:
- Improvement in hepatic inflammation:
- Normalization of ALT:

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Improvement in Knodell HAI inflammation score.

[0062] Sustained loss of serum HCV-RNA correlated with improvement in or resolution of hepatic inflammation. Results demonstrated correlation between sustained virologic response, improvement in hepatic inflammation, normalization of ALT and improvement in HOL.

[0063] The End of Follow-up overall response rate is a composite of the loss of serum HCV-RNA(qPCR) and change in liver histology at end of follow-up (24 weeks following the end of treatment). A patient was classified as an overall responder if HCV-RNA(pPCR) was negative at the 24 week posttreatment evaluation and the posttreatment Knodell HAI inflammation score (sum of categories HHIIII) had improved if the posttreatment value had decreased by 2 or more units relative to the pretreatment score. The percent of usatisand viriolgic responders by time to first negative HCV-RNA, the End of Follow-up (sustained) viriolgic response, histologic response, and overall response rates are summarized in Tables 2, 3, 4 and 5.

End of Follow-up HCV-RNA Sustained Virologic Response: Sustained Loss of HCV-RNA 24 Weeks Following the End of Treatment

[0064] The proportion of patients with eradication of HCV-RNA in the serum 24 weeks following the End of Treatment was two-threefold greater (41% v. 16%) in the group of patients treated with the combination of INTRON® A plus ribavirin compared to those receiving INTRON® A plus placebo.

[0065] The increased length of the combination therapy had the greatest effect on relapser rates. At 24 weeks following End of the Treatment, the relapse rates for 48 weeks of the combination therapy and for 48 weeks of introm A plus placebo were the same (12%). Longer treatment with the combination therapy (48 weeks) and decreased relapse resulted in the highest sustained virologic response rates. Sustained virologic response rates were also significantly higher with 48 weeks of the combination therapy compared to 24 weeks of the combination therapy (38% vs 31%, p value = 0.053.)

10060] Extending the combination therapy from 24 to 48 weeks substantially increased sustained virologic responses in patients who first became HoV-RNA negative at Weeks 12 and 24. The majority of patients who became sustained virologic responders were HCV-RNA negative by Week 4. As summarized in Table 2, the sustained virologic responders were HCV-RNA negative by Week 4. As summarized in Table 2, the sustained virologic responders were substantial portions of the patients on the 48 weeks of combination therapy (HR) at Week 4 of the 48 week treatment and for 81% (3845) of the patients on the 48 weeks of combination therapy at Week 4 of the 48 week treatment. Note that substantial portions of these patients on the 24 and 48 week combination therapy group and 63% of these patients in the 48 week combination therapy group, these responses were sustained. Furthermore, 44% of the patients in the 48 week combination therapy treatment group who first became HCV-RNA negative at Week 24 cheleved a sustained virologic responses. None of the responses that occurred after Week 24 became sustained responders in any treatment group.

[0067] The number of first time responders at Weeks 12 and 24 who become sustained virologic responders 24 weeks after end of treatment were greatest for the patients who received 48 weeks of combination therapy. (See Table 2 below)

Table 2

$Percent \ of \ Sustained \ Virologic \ Responders \ by \ Time \ to \ First \ No \ Detectable \ HCV-RNA \ Levels \ for \ Study \ 1$							
	Intron A	F Ribavirin	Intron A + Placebo				
Time to First No-Detectable HCV-RNA(Weeks)	i						
	24 weeks	48 weeks	24 weeks	48 weeks			
4	81%(35/43)	80%(36/45)	48%(10/21)	70%(14/20)			
12	42%(30/72)	63%(40/63)	9%(3/32)	35%(11/31)			
24	46%(5/11)	44%(11/25)	0%(0/22)	22%(4/18)			

[0068] Table 3 summarizes the End of Follow-up patient response as indicated by serum HCV-RNA.

#### Table 3

Patient Response Status All Treated 95% Confidence Interval	INTRON A	INTRON A + Ribavirin INTRON A		+ Placebo	P-Value <sup>1</sup>	P-Value <sup>1</sup>
	A	В	С	D	B vs D A	A vs D
At EOT	24 weeks	48 weeks	24 weeks	48 weeks		
Negative	121 (53%)	115 (50%)	66 (24%)	54 (24%)	<0.001	<0.001
Positive	107 (47%)	113 (50%)	165 (71%)	171 (76%)		
At End of Follow-up						1
Sustained Responders	70 (31%)	87 (38%)	13 (6%)	29 (13%)	<0.001	<0.001
Relapser	54 (24%)	28 (12%)	53 (23%)	26 (12%)		
Non Responder	104 (46%)	113 (50%)	165 (71%)	170 (76%)	l .	

<sup>&</sup>lt;sup>1</sup>Fisher's Exact Test for End of Treatments P Value A vs B is 0.639; p value for A vs C <0.001.

<sup>2</sup>Logistic Regression for End of Follow-Up Comparisons: P Value A vs B is 0.053; p value for A vs C <0.001.

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[0069] Combination therapy significantly increased the virologic response at the end of treatment compared to IN-TRON A monotherapy. See Table 3. The palues for comparison of the combination therapy for 48 weeks with INTRON A monotherapy for 48 weeks and the comparisons of 24 weeks of combination therapy with 24 and 48 weeks of INTRON A monotherapy are each <0.001. Extending the combination therapy from 24 to 48 weeks decreased the relapse rate by 50% (24% to 12%) thereby making the 48 week combination therapy more effective than the 24 week combination (p=0.053).

[0070] Pre- and Posttreatment biopsies were available for 79% (179/229) and 69% (157/229) of the patients treated with INTRON® A plus trabertin for 24 and 48 weeks respectively and for 76% (156/231) and 70% (158/225) of those patients who received INTRON® A plus placebo for 24 and 48 weeks, respectively. Table 4 summarizes the effect of treatment on hepatic inflammation for patients with both pre- and posttreatment fiver biopsy results. As with the sustained loss of HoV-RNA repictation, the proportion of patients with provement in their inflammation was significantly greater (p<0.001) in patients receiving combination therapy compared to those receiving INTRON® A monotherapy for 48 weeks.

[0071] Eradication of detectable HCV-RNA was highly correlated with normalization of serum ALT. Two to three-fold more patients were ALT normal with the combination therapy compared to INTRON A monotherapy at the End of Foliow-Up. Among patients who had sustained normalized ALT, a higher proportion of 48 week combination therapy patients were sustained virologic responders compared with patients who received 24 weeks of combination therapy and 24 weeks or 48 weeks of INTRON A monotherapy.

# Table 4

End of Follow-up			ver Histology 24 W iell HAI (I+II+III) Sc	eeks Following the ore.	End of Treatm		
Number (%) of Patients <sup>b</sup>							
Patient Status	INTRON A	+ Ribavirin	INTRON A + Placebo		P Value <sup>c</sup>		
	A 24 weeks (N=179) <sup>a</sup>	B 48 weeks (N=157) <sup>a</sup>	C 24 weeks (N+176) <sup>a</sup>	D 48 weeks (N=158) <sup>a</sup>			
Improved Biopsyd	102 (57%)	96 (61%)	77 (44%)	65 (41%)	<0.001		

a N=Number of Patients with paired biopsy samples.

b Patients with both pre- and posttreatment biopsy.

c Fisher's Exact test.

d Change from pretreatment to posttreatment in the Knodell Histological Index (HAI) score (sum of I+II+III) categorized as a decrease of 2 or more from pretreatment.

### Overall Response

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[0072] When the study was designed, it was recognized that because liver biopsy is an invasive procedure that it would be unlikely that postreatment liver biopsies would be obtained for all patients. Therefore, the protocol and statistical analysis plan specified that the analysis for overall response would be based on data for all treated patients and will be estimated by a maximum likelihood method (MLE) for patients whose overall response status could not be determined, i.e., patients with negative HCV-RNA and missing (posttreatment) biopsy evaluations. The protocol also specified that an additional analysis would be performed on patients with both prefreatment and posttreatment biopsy results (i.e., patients with complete data). The overall response is composite of sustained loss of detectable HCV-RNA and improvement in liver histology at End of Follow-Up. Overall response is summarized in Table 5 based on the following analyses:

- maximum likelihood estimate (MLE):
- patients with complete data (results for both pre- and posttreatment biopsy);
- · patients with missing data (either or both HCV/biopsy) treated as failures.

Toble 5

Overall Response Rate.						
Data Analyzed	INTRON A + Ribavirin		INTRON A			
	Α	В	С	D	p value <sup>b</sup>	
	24 weeks	48 weeks	24 weeks	48 weeks	B vs D	
Maximum likelihood estimate <sup>a</sup>	26%	35%	5%	9%	<0.001	
Patients with complete datac	29% (52/179)	41% (64/157)	5% (8/170)	11% (18/158)	<0.001	
Treat missing as failuresd	23% (52/228)	28% (64/228)	28% (64/228)	8% (18/225)	<0.001	

a MLE based on logistic regression. Logistic regression analysis of patients with complete biopsy data The p value for A vs D is <0.001 and for A vs B is 0.096.

vs B is 0.096. b Fisher's exact test.

c Complete data ≈ pre- and posttreatment biopsy .results. (Logistic regression analysis.)

d Patients who had either virology or biopsy data missing or both were counted as non-responders

[0073] As would be anticipated from individual results for effect of treatment on eradication of HCV-RNA at end of follow-up and improvement in hepatic inflammation, the overall response rate in the INTRON® A plus ribavirin 48 week

treatment group was significantly greater (<0.001), than that observed in the INTRON® A plus placebo 48 week treatment group. There was a statistically significant improvement in the overall response with the 48 week combination therapy treatment compared to 48 weeks of INTRON A monotherapy treatment as measured by MLE and complete blopsy. The overall response rates for 24 and 48 weeks of combination therapy treatment was significantly higher than with 24 and 48 weeks of INTRON A monotherapy, respectively.

[0074] Logistic regression analysis was done on all baseline demographic variables and disease characteristics. The only baseline statistically significant patient and disease characteristics predictive of End of Follow-up sustained response were genotype other than 1 and virtu load 25 million.

[0075] For number of viral copies (≤2 million, >2 million), the difference was statistically significant in favor of higher response rates in patients with ≤2 million copies (Table 6 ).

[0078] When genotype and baseline virus load are combined, a hierarchy of response is observed. Those patients with genotype other than 1 and baseline virus load 52, million copies who received 24 and 48 weeks of combination therapy had the best End of Follow-up response; the sustained virologic response for those patients with genotype 1 and >2 million copies who had 48 weeks of the INTROA plus ribavirin combination therapy was two times better than those same type patients who had the combination therapy for only 24 weeks. (See Table 6)

### Table 6

Diaca	oc ontraoteristics vs	Sustained Response		»,		
	Number (%) of Patients					
Disease Characteristic <sup>a</sup>	INTRON A + Ribavirin		INTRON A + Placebo			
	24 weeks	48 weeks	24 weeks	48 weeks		
HCV-RNA/viral load						
≤ 2 million copies/ml	42% (26/62)	43% (33/76)	9% (7/74)	29% (18/63)		
> 2 million copies/mL	27% (44/166)	36% (54/152)	4% (6/157)	7% (11/162)		
HCV Genotype						
1	16 %(26/164)	28% (46/166)	2%(3/167)	7&(11/162)		
Other Genotypes	69% (44/64)	66% (41/62)	16% (10/64)	29% (18/63)		
Genotype/Baseline HCV-RNA						
Other Genotypes ≤2 million copies/ml	88% (14/16)	71 %(15/21)	25% (5/20)	50% (10/20)		
Other Genotypes >2 million copies/ml	63% (30/48)	63% (26/41)	11% (5/44)	19% (8/43)		
Genotype 1, ≤2 million copies/ml	26% (12/46)	33 %(18/55)	4 %(2/54)	19 %(8/43)		
Genotype 1, >2 million copies/ml	12% (14/118)	25 %(28/111)	1% (1/113)	3% (3/119)		

a At entry, patients were stratified by number of HCV viral copies (<2 million, >2 million), genotype (1 or other), and cirrhosis (present or absent) as measured by HCV-RNA/of PCR.

Table 7

Sustained Virologic Response Rates (%) by HCV-Genotype						
	INTRON A + Ribavirin			+ Placebo		
Genotype	24 weeks	48 weeks	24 weeks	48 weeks		
1	16% (27/165)	28% (46/166)	21% (4/168)	7% (11/162)		
2	83% (25/30)	68% (25/37)	18% (7/38)	35% (15/43		
3	57% (16/28)	65% (15/23)	8% (2/24)	16% (3/19)		
4-6	40% (2/5)	50% (1/2)	0	0		

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[0077] Table 7 illustrates that the sustained virologic response rates for patients of each genotype with 48 weeks of the combination therapy were greater than those heated with Intron A plus placebo for 24 and 48 weeks. With the exception of HCV-genotype 2 patients, extending the duration of the combination therapy increased the proportion of patients with sustained virologic responses. But see Table 15 for combined virologic responses for Studies 1 & 2). Logistic regression analysis of sustained virologic response demonstrated that, in addition to treatment group, HCV genotype other than 1 and 52 million copies of baseline HCV virus/mL were significant predictors of sustained virologic response (and with the combination therapy for 48 weeks improved the sustained virologic response rates for patients expected to have the lowest response rates, namely, those patients with HCV-genotype I and greater than 2 million copies of HCV virus/mL. These patients had sustained virologic response rates which were two fold greater than those who had 24 weeks of the virus must be sustained virologic response for HCV-genotype 1 patients who received 48 weeks of combination therapy. Significantly the sustained virologic response for HCV-genotype 1 patients who received 48 weeks of combination therapy was 1.75 times greater than that for those who received 42 weeks of the virus from the rate of the virus of virus of the virus of the virus of virus of

#### Study 2:

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[0078] By basically the same methodology as described above in Study 1, Study 2 was also conducted in 43 international sites (322 patients) using the following three treatment regimens:
[0079] The results are summarized below:

- INTRON® A 3 million IU SC TIW plus ribavirin 1000 or 1200 mg/day PO in two divided doses for 24 weeks: or
- INTRON® A 3 million IU SC TIW plus ribavirin 1000 or 1200 mg/day
   PO in two divided doses for 48 weeks: or
- 3. INTRON® A 3 million IU SC TIW plus placebo matching ribavirin
- 25 PO in two divided doses for 48 weeks

# Efficacy

[0080] The primary efficacy objective is sustained virologic response as defined by the loss of detectable serum HCV-RNA(qPCR) measured at End of Follow-up (24 weeks following the end of treatment). A patient was classified as an overall responder if HCV-RNA(PCR) was negative at the 24 week posttreatment evaluation and the posttreatment Knodel HAI inflammation score (sum of categories HiHIII) had improved (decreased) by 2 or more units relative to the pretreatment score. The percent of sustained virologic responders by time to first negative HCV-RNA, the End of Follow-up virologic response, histologic response, and overall response rates are summarized in Tables 9, 10,11 and 12.

[0081] Patient body weight and their baseline disease characteristics (HCV genotype and initial viral load) for all patients in Study 2 is given in Table 8 below.

# Table 8.

	INTRON A	+ Ribavirin	INTRON A + Placel	
Body Weight	24 weeks	48 weeks	48 weeks	
>75 kg	116 (42%)	129 (47%)	122 (44%)	
≤ <b>7</b> 5 kg	161 (58%)	148 (53%)	156 (56%)	
CV Genotype				
1	161 (58%)	159 (57%)	168 (60%)	
2	27 (10%)	23 (8%)	21 (8%)	
3	73 (26%)	74 (27%)	79 (28%)	
4	12 (4%)	16 (6%)	9 (3%)	
5	1 (0.4%)	5 (2%)	1 (0.4%)	
6	3 (1%)	0	0	

# Table 8. (continued)

Body Weight and Baseline Disease Characteristics for Study 2 Patients							
	INTRON A	+ Ribavirin	INTRON A + Placebo				
HCV-RNA/of PCR (copies/ml)							
mean geometric	2,229,797	2,064,959	2,351,824				
≤ 2 million copies/mL	108 (39%)	115 (42%)	95 (34%)				
>2 million copies/mL	169 (61%)	162 (59%)	183 (66%)				

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Tab	e 9.				
Percent of Sustained Virologic Responders by Time	to First No Dete	ctable HCV-RN	A Levels for Study		
Time to First No-Detectable HCV-RNA (Weeks) Intron A + Ribavirin Intron A+					
	24 weeks	48 weeks	48 weeks		
4	84%(57/68)	83%(58/70)	72%(33/46)		
12	47%(36/77)	69%(51/74)	34%(18/53)		
24	12%(3/26)	45%(9/20)	10%(2/21)		

[0082] As summarized in Table 9, the majority of patients who became sustained virdogic responders had negative HCV-RNA levels by Week 4 of treatment. However, substantial proportions of patients in the 24 and 48 week combination therapy treatment groups who were HCV-RNA positive at Week 4 responded for the first time at Week 12; 47% of those in the 24 Week treatment group became sustained responders. Importantly, 45% (9/20) of the patients in the 48 Week treatment group became sustained responders. Importantly, 45% (9/20) of the patients in the 48 Week combination therapy treatment group who first became negative at Week 24 became sustained virologic responders. None of the responses that occurred after Week 24 became sustained in any of the three treatment groups.

# End of Follow-up HCV-RNA Response: Sustained Loss of HCV-RNA 24 Weeks Following the End of Treatment

[0083] The proportion of patients with eradication of HCV-RNA in the serum 24 weeks following the end of the combination therapy treatment was significantly greater in patients treated with the combination therapy of INTRON® A Pulsa ribavirin compared to those receiving INTRON® A monotherapy. Table 10 summarizes the End of Follow-up patient response as indicated by serum HCV-RNA.

Table 10.

	Number (%) of Patients						
	INTRON A	+ Ribavirin	INTRON A + Placebo				
	A	В	С	p value			
	24 weeks (N=277)	48 weeks (N=277)	48 weeks (N=278)	B vs C1			
EOT <sup>2</sup>							
Negative	57% (157)	52% (145)	33% (93)	<0.001			
Positive	42% (120)	42% (132)	65% (185)				
EOFU <sup>3</sup>	L		L				
Sustained	35% (96)	43% (118)	19% (53)	<0.001			
Relapse	23% (23)	10% (27)	15% (41)				
Non Responders	42% (42)	48% (132)	66% (184)				

<sup>&</sup>lt;sup>1</sup>Fisher's Exact Test for EOT comparisons; logistic regression for EOFU.

<sup>2</sup>EOT = End of Treatment. P value for A vs C = <0.001 and for A vs B is 0.348.

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[0084] Table 10 illustrates that (1) the 24 and 48 week combination therapies significantly increased the sustained virologic response at the end of treatment compared to that for the intron A monotherapy (p values are both <0.001) and (2) the increasing the length of the Combination Therapy from 24 to 48 weeks had the greatest effect on relapse rates (10% for 48 weeks vs 23% for 24 weeks, p value is 0.055.)

End of Follow-up Liver Histology: Improvement in Liver Histology 24 Weeks Following the End of Treatment Based on Knodell Histological Activity Index (HAI) Scores (I+II+III)

[0085] Pre- and Posttreatment biopsies were available for 74% (204/277) and 60% (167/277) of the patients treated with INTRON® A plus ribavirin for 24 and 48 weeks, respectively, and for 69% (191/278) of those patients who received INTRON® A plus placebot. Table 11 summarizes the effect of treatment on hepatic inflammation for patients with both pre- and posttreatment liver biopsy results. As with the sustained loss of HCV-RNA replication, the proportion of patients with improvement in liver inflammation was significantly greater (p-0.001) in patients receiving combination therapy for 48 weeks compared to those receiving INTRON® A monotherapy for 48 weeks. Extending the combinatin therapy from 24 to 48 weeks also significantly increased the proportion of patients who had improvement in hepatic inflammation (p value=0.046).

<sup>3</sup>EOFU = End of Follow-Up. P value for A vs C = <0.001 and for A vs B is 0.348.

#### Table 11.

End of Folbw-up Live		ement in Liver Histolo n the Knodell HAI (I+I	gy 24 Weeks Following I+III) Score.	the End of Treatmen			
T	Number (%) of Patients <sup>b</sup>						
Ī	INTRON A	+ Ribavirin	INTRON A + Placebo				
Patient Status	Α	В	С	p value <sup>c</sup>			
	24 weeks (N=204) <sup>a</sup>	48 weeks (N=167) <sup>a</sup>	48 weeks (N=191) <sup>a</sup>	B vs C			
Improved Biopsy <sup>d</sup>	53% (107)	63% (105)	39% (74)	<0.001			

a Number of patients wit paired biopsies.

#### Overall Response

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[0086] The overall response is summarized in Table 12 based on the following analyses:

- maximum likelihood estimate (MLE);
- patients with complete data (results for both pre- and posttreatment biopsy);
  - patients with missing data (either or both HCV-RNA/biopsy) treated as failures.

Table 12.

Ove	rall Response Ra	ite.		
	INTRON A	INTRON A + Placebo		
Data Analyzed	Α	В	С	
	24 weeks	48 weeks	48 weeks	
ML Estimate	28%	37%	17%	
Patients with Complete Biopsy Datac	30% (62/204)	41% (68/167)	17% (32/191)	
Treat Missing as Failuresd	22% (62/277)	24% (68/277)	12% (32/278)	

a MIE based on logistic regression. P value for B vs C = <0.001 and for A vs C = <0.002 and for A vs B is 0.043.

[0087] As would be anticipated from individual results for effect of treatment on eradication of HCV-RNA at Importance of Follow-up and improvement in hepatic inflammation, the overall response rate in the INTRON A plus having may be significantly greater with a two fold improvement over that observed with INTRON A plus placebo groups for all methods of evaluation.

[0088] Logistic regression analysis was done on all baseline demographic variables and disease characteristics. The only baseline statistically significant characteristic predictive of End of Follow-up sustained response was genotype other than 1.

Table 13.

	Intron A	Ribavirin	Intron A + Placebo
Genotype	24 Weeks	48 Weeks	48 Week
1	18%(29/161)	30% (48/159)	11% (19/168)
2	59% (16/27)	74% (17/23)	35% (8/23)

b Patients with both pre-and posttreatment biopsy.

c Fisher's Exact test. P value for A vs C = 0.007 and A for A vs B is 0.046,

d Change from pretreatment to posttreatment in the Knodell Histological Index (HAI) score (sum of I+II+III) categorized as a decrease of 2 or more from pretreatment.

b Fisher's Exact test

c Complete data = pre- and posttreatment biopsy results. P value for B vs C is <0.001, for A vs C is <0.005 and for A vs B is 0.032.

d Patients who had either virology or biopsy data missing or both were counted as failures...

Table 13. (continued)

Sustained Viro	logic Response Ra	ates by HCV Genot	ype for Study 2 Patients
	Intron A + Ribavirin		Intron A + Placebo
Genotype	24 Weeks	48 Weeks	48 Week
3	66% (48/73)	61% (45/74)	32% (25/79)
4-6	19% (3/16)	38% (8/21)	12%(1/8)

[0089] The combination therapy provided higher sustained virologic response rate for all genotype compared to Intron A plus Placebo. Extending the duration of the combination therapy to 48 weeks increased the proportion of sustained virologic response for all genotypes except type 3. (See Table 13 as well as Table 15 for the combined vivologic response for studies 1 and 2)

[0090] For number of viral copies (≤2 million, >2 million), there was a numerical difference in favor of higher sustained virologic response rates in patients with ≤2 million copies (Table 14). When genotype and baseline virus load are combined, a hierarchy of response is observed. Those patients with genotype other than 1 and baseline virus load ≤2 million copies had the best End of Follow-Up response; those patients with genotype 1 and >2 million copies who received the longer 4 week treatment with combination therapy had the most significant improvement in substantial virologic response of all the groups.

Table 14

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	Table 14				
Disease Characteristics vs Sustained Viro	ologic Response	: All-Treated Pat	ients In Study No. 2		
	umber (%) of Patients				
Disease Characteristic	INTRON® A	plus ribavirin	INTRON® A plus Placel		
	24 weeks	48 weeks	48 weeks		
HCV-RNA/qPCR					
≤2 million copies/mL	44% (48/108)	47% (54/115)	31 % (15/49)		
> 2 million copies/mL	28% (48/109)	40% (64/102)	13% (24/183)		
HCV Genotype					
1	18% (29/161)	30% (48/159)	11% (19/168)		
Other Genotype	58% (67/116)	59% (70/118)	31% (34/110)		
Genotype/Baseline HCV-RNA/qPCR	l				
Other genotype /≤2 million copies/mL	53% (28/53)	62% (34/55)	31% (15/49)		
Other genotype /> 2 million copies/mL	62% (39/63)	57% (36/63)	31% (19/61)		
Genotype1&≤2 million copies/mL	36% (20/55)	33% (20/60)	30% (14/46)		
Genotype1&> 2 million copies/mL	8% (9/106)	28% (28/99)	4% (5/122)		

[0091] Table 14 illustrates that extending the combination therapy to 48 weeks generally improved the sustained virologic response rates. The highest sustained virologic response rates were observed with patients who received the combination therapy for 48 weeks with genotypes other than 1 and initial HCV levels of 22 million copiesmL importantly, for patients with genotype 1 and HCV levels >2 million, sustained virologic response rates with 48 weeks of combination therapy were more than 3 times higher than rates with only 24 week of the combination.

# Conclusions on Efficacy for Combined Results for All Patients in Studies 1 + 2

[0092] Combination therapy was significantly more efficacious than INTRON®A monotherapy for initial treatment of chronic hepatitis C. Sustained virologic response rates in these antiviral treatment naive patients were almost three

times higher with 48 weeks of combination therapy compared with 48 weeks of Intron A monotherapy and significantly higher with 48 weeks compared to 24 weeks of the combination therapy. The improvement in sustained response rates could be accounted for by two treatment effects: higher response rates at the end of treatment and decreased relapse rates. The net result of both of these effects was the highest sustained response rate with the 48 weeks of combination therapy compared to 48 weeks of monotherapy or a shorter regimen with the combination. This enhancement of efficacy with 48 weeks of combination therapy also included other measures of response such as biochemical (ALT) and histological endopriots.

[0093] In fact, sustained loss of serum HCV-RNA was highly correlated with other clinical endpoints - normalization of ALT and improvement in or resolution of hepatic inflammation. Loss of detectable HCV-RNA at the end of follow-up was associated with normalization of ALT in all treatment groups but was somewhat higher with combination therapy. The majority of combination therapy patients who were ALT normal were also HCV-RNA negative (83-87%).

[0094] Increased length of therapy had the greatest effect on relapser rates. At End of Follow Up, the relapse rates for both the 48 week combination and monotherapy treatment groups were lower than the relapse rate of the 24 week combination therapy treatment group. The combination of the high end of treatment response resulting from 24 and 48 weeks of the combination therapy compared to 48 weeks of lintron A monotherapy and decreased relapse rates resulted in the highest sustained response rates. Sustained responses rates were twice as high with 48 weeks of combination therapy compared with Intron A monotherapy(49) (p<0.001). Sustained virologic response rates were also higher with 48 weeks of combination therapy compared to nity 24 weeks of the Combination the nity 24 weeks of the Combination therapy compared to

[0095] The benefit of combination therapy was maintained irrespective of standard predictors of response to IN-TRONDAM montherapy and 24-week combination therapy in relapsed patients. At entry to these trials, patients were stratified by the following disease characteristics: HCV genotype (type 1 or other genotypes); extent of HCV virus level (number of virus copies in the serum 22 million/III. or 22 million/III.); and cirricots (greesn) or absent), Logistic regression analysis of sustained virologic response demonstrated that, in addition to treatment group, only HCV genotype was a significant predictor of sustained virologic response. Neither pretreatment HCV virus level no presence of cirricots appeared to influence the ability of previously untreated patients to achieve sustained virologic response to the combination therapy.

[0096] With 48 weeks of combination therapy, sustained response rates were consistently higher than those for 48 weeks of Intron A monotherapy regardless of generopye and were generally higher than 24 weeks of combination therapy. As a group, patients infected with genotypes 1 have been shown to be less responsive to INTRON®A monotherapy than patients infected with other genotypes. Despite this, sustained virologic response rates for genotype 1 were approximately 3 times higher with 48 weeks of combination therapy compared to 48 weeks of Intron A monotherapy and almost twice as high with 48 weeks compared to 24 weeks of combination therapy. The combination therapy consistently produced higher sustained virologic response rates in patients infected with other genotypes. Response rates with both 24 and 48 weeks of combination therapy were higher than with 48 weeks of Intron A monotherapy and in all genotypes except types 2.8 3 which genotypes had the same response rate Le, about 46% at the end of 24 weeks as well as after 48 weeks of combination therapy), extending the duration of the combination therapy to 48 weeks increased the proportion of patients with sustained virologic responses. (See Table 15)

[0097] The combination therapy was also more effective than 48 weeks of Intron A monotherapy in producing sustained virologic responses regardless of virus level at Baseline. Forty-eight weeks of combination therapy consistently produced higher sustained virologic response rates at every level of virus infection than 48 weeks of Intron A monotherapy. Sustained response rates were similar for most virus levels in both the 24 and 48 week combination therapy treatment groups, except with the highest virus levels (5–6 and 240Xf0 copies/ml), where sustained response rates with the 48 week combination therapy groups were approximately twice as high as the 24 week combination therapy groups were approximately twice as high as the 24 week combination therapy groups.

fo [0093] As noted previously, patients with genotypes other than type 1 had higher sustained virologic response rates than those with type 1 and a virus level 25 million/mt.. as associated with a better response rate than 1-2 million/mt.. It was notable that extending the combination therapy treatment to 43 weeks improved the sustained response rates for patients expected to have the lowest response rates, namely, those with genotype 1 and 2-2 million copies of HCV-RNA/mt. Extending the combinatin therapy to 48 weeks in this group of patients produced sustained virologic response rates that were 4 times higher than those with only 24 weeks of combination therapy.

[0099] Other demographic/disease history characteristics had little effect on outcome with combination therapy, in contrast, considerably lower sustained response rates were noted with 48 weeks of intron A monotherapy in patients older than 55 years, >75 kg or who were infected through transfusion. All had sustained response rates in the range of 10-12%.

[0100] Availability of paired biopsies was high compared with similar types of studies in chronic hepatitis C patients. As anticipated, pre- and/or postfreatment liver biopsies were unavailable for a proportion of patients for a variety of reasons. However, paired biopsies were obtained from 71% of patients. Improvement was noted in a significantly higher proportion of 48 week combinatin therapy patients compared with the 48 week futron A monotherapy patients at End.

of Follow Up (p<0.001). Twenty-four weeks of combinatin therapy also was significantly more efficacious than 48 weeks of Intron A monotherapy in improving hepatic inflammation. As noted previously, the correlation with virologic response was maintained whether biopsies were assessed by improvement or mean change from baseline in necroillammation; score; 64-69% of patients had improvement in hepatic inflammation with mean changes from baseline of -3.8 to -5.0. The most substantial mean change was in the 48 week Intron A monotherapy patients.

[0101] As anticipated, sustained vivologic responders in all treatment groups experienced greater improvement in liver biopsy inflammation scores than patients who remained HCV-RNA positive, and although the extent of improvement was similar in all groups the proportion of vivologic responders with histologic improvement was a fleast twice a high with both the 24 and 48 weeks of combination therapy than with 48 weeks of Intron A monotherapy. Extending the combination therapy resulted in higher mean improvements in hepatic inflammation. It is also interesting to note that patients who relapsed from the 48 week combination therapy had considerable mean improvement in inflammation. The combined results are summarized in Tables 15 to 20.

Table 15. Combined Virologic Response for Studies 1 and

Efficacy: Loss of Delectable HCV-RNA at End of Treatment (EOT and Sustained Virologic Response At End of Follow Up (EOFU) and Percent Patients Responding

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	Intron A + Ribavirin Intron A + Placebo		+ Placebo	P Value		
	24	48	24	48	l	
	weeks	weeks	weeks	weeks	L	
No. of	N=505	N=505	N=231	N=503	B vs D	A vs D
Patients <sup>1</sup>	A	В	С	D		
EOT <sup>2</sup>	53%	50%	29%	24%	<0.001	<0.001
EOFU <sup>3</sup>	33%±4	41%±5	6%	16%	<0.001	<0.001
HCV	(No. of	(No. of				
Genotype	Patients)	Patients)				
Type 1	· 17%±4	29%±5	2%	9%		
	(56/326)	(94/325)				
2	73%±124	70%±12⁴	18%	35%	j	
	(41/57)	(42/60)			}	
3	63%±94	62%±104	8%	29%		
1	(64/101)	(60/97)		ļ		
Types	24%±10	39%±20	0%	11%	- 1	
4/5/6	(5/21)	(9/23)				

Patient characteristics: (a) 66%male; 34% female; (b) mean age: 42.7 yrs; (c) Mean pretreatment HCV-RNA levels ~2 million copies/mi: 34%, and >2 million copies/mi: 66%; Super Quant, NGI; (d) HCV genotype 1=66%; genotype 2=15%, non 1-3-35%.

EOT is End of Treatment. EOFU is End of Follow-Up - 24 weeks posttreatment; P value A vs B≈ 0.257;

Sustained virologic response at End of Follow-Up; P Value A vs B = 0.008

<sup>4</sup> The sustained virologic response for HCV genotypes 2 and 3 at the end of 24 weeks was about 64.5% and at the end of 48 weeks was about 65%.

Table 16

Percentage of Sustained Virologic R	Intron A		A + Placebo	
Patient	24 weeks	48 weeks	24 weeks	48 weeks
HCV-Genotype 1 and ≤2 million copies/mL	32%	33%	4%	25%
HCV Genotype 1 and >2 million copies/mL	10%	27%	1%	33%
All HCV-Genotypes 1	17%	29%	2%	9%

[0102] The sustained virologic response for all HCV genotype 1 patients treated for 24 and 48 weeks with the Combination Therapy was statistically significantly superior to that observed for all HCV genotype I patients treated with INTRON A plus placebo for 24 and 48 weeks. The sustained virologic response for HCV-genotype 1 patients with a baseline HCV viral load of greater than 2 million copies/mL was statistically significantly superior at 24 and 48 weeks of Combination Therapy compared to INTRON A + Placebo for 24 and 48 weeks.

Table 17.

	Intron A + Ribavirin		Intron A	Placebo
Patients				
	24 weeks	48 weeks	24 weeks	48 weeks
HCV Genotype 1 and Baseline HCV-RNA levels of ≤ 2 million copies/mL	32%	33%	4%	25%
HCV Genotype 1 and Baseline HCV-RNA levels of > 2 million copies/mL	11 %	27%	1 %	4%
Other HCV Genotypes <sup>1</sup> and Baseline HCV-RNA levels of > 2 million	61 %	64%	25%	36%
Other HCV Genotypes <sup>1</sup> and Baseline HCV-RNA levels of > 2 million copies	62%	61 %	11 %	10%

<sup>1</sup> HCV-genotypes 4/5/6 account for 3% of patients Sustained Response for I+R: 24% (24 wks) and 38% (48 weeks) for I+P: 0% (24 weeks) end 11 % (48 weeks) Sustained Response Rate for HCV-Genotypes 2+3 is same as above

Table 18.

Sustained Virologic Response by Baseline HCV-RNA Levels for ALL Patients (Studies 1 & 2)							
Patients	Intron A	Intron A	Intron A + Placebo				
	24 weeks	48 weeks	24 weeks	48 weeks			
Patients with Baseline HCV-RNA Levels ≤2 million copies/mL	44%	46%	9%	36%			
Patients with Baseline HCV-RNA Levels > 2 million copies/mL	27%	38%	4%	10%			

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Table 10

	Table 10.						
Percent of Sustained Virologic Responders by Time to First No Detectable HCV-RNA Levels for Studies 1 & 2							
	Intron A	Ribavirin	Intron A + Placebo				
Time to First No-Detectable HCV-RNA (Weeks)	First No-Detectable HCV-RNA (Weeks)						
	24 weeks	48 weeks	24 weeks	48 weeks			
4	83%(92/111)	82%(94/115)	48%(10/21)	71%(47/66			
12	44%(66/149)	66%(91/137)	9%(3/32)	35%(29/84			
24	19%(8/42)	44%(20/45)	0%(0/22)	15%(6/39)			

Table 20.

			Table 20.			
	ALT Response:	Normalization of	Serum ALT in Al	Patients from S	tudies 1 & 2	
	INTRON A	+ Ribavirin	INTRON A	+ Placebo		
At End of Treatment	· A	В	С	D	P <sup>a</sup> Value	Pª Value
	24 weeks (N=505)	48 weeks (N=505)	24 weeks (N=231)	48 weeks (N=503)	B vs D	A vs D
Study						
1 & 2 <sup>b</sup>	66% (329)	66% (334)	24% (56)	37% (185)	<0.001	0.739
1	58% (133)	61% (138)	24% (56)	28% (62)	<0.001	<0.001
2	71% (196)	71% (196)	-	44% (123)	<0.001	<0.001
End of Follow	- Up Study		L			1
1+2 <sup>b</sup>	36% (181)	44% (221)	11% (25)	24% (102)	<0.001	<0.001
1	32% (72)	36% (83)	11% (25)	16% (35)	<0.001	<0.001
2	39% (109)	50% (138)	-	24% (67)	<0.001	<0.001

<sup>&</sup>lt;sup>8</sup> Cochran-Mantet Haenszei general association for combined results; Fisher's Exact Test for individual studies.
<sup>b</sup> Combined results

#### Claims

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- 1. The use of ribavirin for the manufacture of a pharmaceutical composition for treating a patient having chronic hepatitis C infection to eradicate detectable HCV-RNA wherein the pharmaceutical composition is for adminishering an effective amount of interform alpha, characterised in that the ribavirin in association with the interferon alpha is for administration for a time period of 40-50 weeks, the patient is an antiviral treatment naive patient, and the patient is one having a HCV gendropke pute 1 infection and a viral load of greater than 2 million copies per mill of serum as measured by HCV-RNA quantitative PCR.
- 2. The use of interferon alpha for the manufacture of a pharmaceutical composition for treating a patient having chronic hepatitis C infection to eradicate detectable HCV-RNA wherein the pharmaceutical composition is for administering an effective amount of interferon alpha in association with an effective amount of ribavirin, characterised in that the ribavirin in association with the interferon alpha is or administration for a time period of 40-50 weeks, the patient is an antiviral treatment naive patient, and the patient is one having a HCV genotype type 1 infection and a viral load of greater than 2 million copies per ml of serum as measured by HCV-RNA quantitative PCR.
  - 3. The use of both ribavirin and interferon alpha for the manufacture of pharmaceutical compositions for treating a

patient having chronic hepatitis C infection to eradicate detectable HCV-RNA wherein the pharmacoutical composition is for administering an effective amount of ribavinin in association with an effective amount of interferon alpha is characterised in that the ribavinin in association with the interferon alpha is for administration for a time period of 46-50 weeks, the patient is an antiviral treatment nave patient, and the patient is one having a HCV genotype type 1 infection and a viral load of greater than 2 million copies per ml of serum as measured by HCV-RNA quantitative PCR.

- 4. The use as claimed in any one of Claims 1 to 3 wherein the interferon-alpha administered is selected from interferon alpha-2a, interferon alpha-2b, a consensus interferon, a purified interferon alpha product, a pegylated interferon alpha-2b.
- 5. The use according to any one of the preceding claims wherein the interferon alpha is a pegylated interferon alpha.
- 6. The use according to Claim 5 wherein the interferon is pegylated interferon alpha-2b.
- The use according to Claim 5 wherein the interferon is pegylated interferon alpha-2a.
- 8. The use as claimed in any one of Claims 1 to 4 wherein the interferon alpha is interferon alpha-2b.
- The use as claimed in any one of claims 1 to 4 wherein the interferon employed is interferon alpha-2a.
  - 10. The use as claimed in any one of the preceding claims wherein the amount of ribavirin to be administered is 800 to 1200 mg per day.
- 11. The use as claimed in any one of the Claims 1 to 10 wherein the amount of alpha interferon to be administered is from 2 to 10 million IU per week on a weekly. TIW, QOD, or daily basis.

# Patentansprüche

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- 1. Verwendung von Ribavirin zur Herstellung einer pharmazeutischen Zusammensetzung zur Behandlung eines Patienten mit knonischer Hepstellt Sc Infektion, um nachweisbere HÜV-RNA auszumezen, wobei die pharmazeutische Zusammensetzung zur Verabreichung einer wirksamen Menge Ribavirin in Verbindung mit einer wirksamen Menge Interferon-cu vorgesehen ist, dadurch gekennzeichent, dass das Ribavirin in Verbindung mit dem Interferon-a zur Verabreichung über einen Zeitraum von 40 bis 50 Wochen vorgesehen ist, der Patient ein im Hinblick auf antivirale Behandlung nahre Patient ist und der Patient ein Infektion mit einem HCV-Genoty Typ 1 und eine virale Belastung von mehr als 2 Million Kopien/ml Serum, wie mittels HCV-RNA quantitativer PCR bestimmt, aufwelst.
- Verwendung von Interferon-α zur Herstellung einer pharmazeutischen Zusammensetzung zur Behandlung eines
  Patienten mit chronischer Hepatikis C Infektion, um nachwielsbare HCV-RNA auszumerzen, wobei die pharmazeutische Zusammensetzung zur Verabreichung einer wirksamen Meng enterferon-α in Verbindung mit einer wirksamen Menge Ribavirin vorgesehen ist, dadurch gekennzeichnet, dass das Ribavirin in Verbindung mit dem
  Interferon-α zur Verabreichung für einen Zeitraum von 40 bis 50 Wochen vorgesehen ist. Per Zeitent im Hinblück
  auf antivirale Behandlung ein nalver Patient ist und der Patient eine Infektion mit HCV-Genotyp Typ 1 und eine
  virale Belastung von mehr als 2 Million Kopien/ml Serum, wie mittels HCV-RNA quantitativer PCR bestimmt, aufweist.
- 3. Verwendung sowohl von Ribawirin als auch Interferon-α für die Herstellung einer pharmazudischen Zusammensetzung zur Behandlung eines Patienten mit chronischer Hepatitis C Infektion, um nachweisbare HCV-RNA auszumerzen, wobei die pharmazeutische Zusammensetzung zur Verabreichung einer wirksamen Menge Ribavirin in Verbindung mit einer wirksamen Menge Interferon-a vorgesehen ist, dadurch gekennzeichnet, dass das Ribavirin in Verbindung mit dem Interferon-a zur Verabreichung über einen Zeitraum von 40 bis 50 Wochen vorgesehen ist, der Patient ein im Hinblick auf antivirale Behandlung naiver Patient ist und der Patient eine Infektion des HCV-Genochy Typ 1 und eine virale Belastung von mehr als 2 Million Kopien/ml Serum, wie mittels HCV-RNA quantitativer PCR bestämmt, aufweist.
  - 4. Verwendung gemäß irgendeinem der Ansprüche 1 bis 3, bei der das verabreichte Interferon-α aus Interferon-α-

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- 2a, Interferon-α-2b, Consensus-Interferon, gereinigtem Interferon-α-Produkt, pegyliertem Interferon-α-2a oder pegyliertem Interferon-α-2b ausgewählt wird.
- Verwendung gemäß irgendeinem der vorstehenden Ansprüche, bei der das Interferon-α ein pegyliertes Interferonα ist.
- 6. Verwendung gemäß Anspruch 5, bei der das Interferon pegyliertes Interferon-α-2b ist.
- 7. Verwendung gemäß Anspruch 5, bei der das Interferon pegyliertes Interferon-α-2a ist.
- Verwendung gemäß irgendeinem der Ansprüche 1 bis 4, bei der das Interferon-α Interferon-α-2b ist.
- 9. Verwendung gemäß irgendeinem der Ansprüche 1 bis 4, bei der das verwendete Interferon-α Interferon-α-2a ist.
- 10. Verwendung gemäß irgendeinem der vorstehenden Ansprüche, bei der die Menge des verabreichten Ribavirin 800 bis 1200 mc/Tag beträgt.
  - Verwendung gemäß irgendeinem der Ansprüche 1 bis 10, bei der die Menge des verabreichten Interferon-α von 2 bis 10 Millionen IU pro Woche auf einer wöchentlichen TIW, QOD oder täglichen Basis ist.

# Revendications

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- 1. Utilisation de ribavirine pour la fabrication d'une composition pharmaceutique pour traitor un patient souffrant d'une infection par hépatite C chronique, pour éractiquer FARM désetable du VHCI, a composition pharmaceutique étant destinée à administrer une quantité efficace de ribavirine en association avec une quantité efficace d'interféron aipha, ceractérisée en ce que la ribavirine en association avec l'interféron alpha est destinée à têtre administrée pendant une durée de 40 50 semaines, le patient déat un patient natifiérat et le patient étant un patient souffrant d'une infection par le VHC de génotype de type 1 avec une charge virale, mesurée par PCR quantitativé de fARM du VHCs, supérieure à 2 millions de copies par m de sérum.
- 2. Utilisation d'interféron alpha pour la fabrication d'une composition pharmaceutique pour traiter un patient souffrant d'une infection par hépatite C chronique, pour réadiquer /ARN détectable du Vinc, la composition pharmaceutique étant destinée à administrer une quantité efficace d'interféron alpha en association avec une quantité efficace de ribavirine, caractérisée en ce que la ribavirine en association avec finterféron alpha est destinée à être administrée pendant une durée de 40 -50 semaines, le patient étant un patient natifival et le patient étant un patient natifival et le patient étant un patient souffrant d'une infection par le VHC de génotype de type 1 avec une charge virale, mesurée par PCR quantitative de fARNI du VIR, supérieure à z millions de copies par mt de sérvice.
- 40 3. Utilisation conjointe de ribavirine et d'interféron alpha pour la fabrication de compositions pharmaceutiques pour traiter un patient souffrant d'une infection par hépatite C chronique, pour réadiquer I/RAN détectable du VHC, la composition pharmaceutique étant destinée à administrer une quantité efficace d'interféron alpha en association avec une quantité efficace de ribavirine, caractérisée en ce que la ribavirine en association est destinée à être administrée pendant une durée de 40 50 semaines, le patient étant un patient native à-vis d'un traitement antiviral et le patient étant un patient souffrant d'une infection par le VHC de génotype de type 1 avec une charge virale, mesurée par PCR quantitative de l'ARN du VHC, supérieure à 2 millions de copies par mi de sérum.
- Utilisation selon l'une quelconque des revendications 1 à 3, dans laquelle l'interféron alpha administré est sélectionné parmi l'interféron alpha 2a, l'interféron alpha 2, pur linterféron consensus, un produit punifié d'interféron alpha, un interféron alpha 2 polyéthyène glycolé ou un interféron alpha 2b polyéthyène glycolé.
  - Utilisation selon l'une quelconque des revendications précédentes, dans laquelle l'interféron alpha est un interféron alpha polyéthylène glycolé.
  - 6. Utilisation selon la revendication 5, dans laquelle l'interféron alpha est un interféron alpha 2b polyéthylène glycolé.
  - 7. Utilisation selon la revendication 5, dans laquelle l'interféron alpha est un interféron alpha 2a polyéthylène glycolé.

- 8. Utilisation selon l'une quelconque des revendications 1 à 4, dans laquelle l'interféron alpha est l'interféron alpha 2b.
- 9. Utilisation selon l'une quelconque des revendications 1 à 4, dans laquelle l'interféron alpha est l'interféron alpha 2a.
- Utilisation selon l'une quelconque des revendications précédentes, dans laquelle la quantité de ribavirine à administrer est de 800 à 1 200 mg par jour.
  - 11. Utilisation selon l'une quelconque des revendications 1 à 10, dans laquelle la quantité d'interféron alpha à administrer est de 2 à 10 millions Ul par semaine sur base d'une dose hebdomadaire, de deux doses par semaine (TIVN), d'une dose tous les deux jours (2000) ou d'une dose journalière.

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